

MARKSCHEME

May 2006

CHEMISTRY

Higher Level

Paper 2

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SECTION A

1. (a) $\text{C}_6\text{H}_{12} + 9\text{O}_2 \rightarrow 6\text{CO}_2 + 6\text{H}_2\text{O};$ [1]
- (b) (i) $(\Delta H^\ominus = \sum \Delta H_{\text{f products}}^\ominus - \sum \Delta H_{\text{f reactants}}^\ominus)$
 $\Delta H^\ominus = (6 \times -394 + 6 \times -242) - (-43);$
 $\Delta H_c^\ominus = -3773 / -3.8 \times 10^3 \text{ (kJ mol}^{-1}\text{);}$ [2]
Accept 2, 3 or 4 sf.
Award [1] for $+3773 / +3.8 \times 10^3 \text{ (kJ mol}^{-1}\text{)}.$
Allow ECF from (a) only if coefficients used.
- (ii) $\Delta S^\ominus = (S_{\text{p}}^\ominus - S_{\text{r}}^\ominus) = (6 \times 189 + 6 \times 214) - (385 + 9 \times 205);$
 $\Delta S_c^\ominus = 188 \text{ (J K}^{-1} \text{ mol}^{-1}\text{);}$ [2]
Accept only 3sf.
Award [1] for $-188.$
Allow ECF from (a) only if coefficients used.
- (c) $(\Delta G_c^\ominus = \Delta H_c^\ominus - T\Delta S_c^\ominus) = -3800 - (298 \times 0.188);$
 $= -3900 \text{ kJ mol}^{-1}.$ [2]
Accept -3800 to $-3900.$
Accept 2, 3 or 4 sf.
Allow ECF from (b).
Units needed for second mark.
- (d) spontaneous and ΔG^\ominus negative; [1]
Allow ECF from (c).
- (e) $-1 \times \Delta H_1 / 676;$
 $1 \times \Delta H_2 / -394;$
 $2 \times \Delta H_3 / -484;$
 $\Delta H_4 = -202 \text{ (kJ mol}^{-1}\text{);}$ [4]
Accept alternative methods.
Correct answers score [4].
Award [3] for $(+)202$ or $(+)40 \text{ (kJ / kJ mol}^{-1}\text{)}.$

2. (a) $A_r(\text{Tl}) = 203 \times 0.2952 + 205 \times 0.7048 / A_r(\text{Tl}) = 204.41$;
 $A_r(\text{Br}) = 79 \times 0.5069 + 81 \times 0.4931 / A_r(\text{Br}) = 79.99$;
 $M_r(\text{TlBr}_3) = 204.41 + 3 \times 79.99 = 444.38 / 444.37$; [3]
 Correct answer scores [3].
 Ignore units of g or g mol^{-1} .
 Apply ECF to M_r from A_r values.
- (b) M_r is an average value (because of the isotopes);
 each HBr molecule has its own value depending on which isotopes (of H or Br) it contains/OWTTE; [2]
- (c) $1s^2 2s^2 2p^6 3s^2 3p^6 3d^{10} 4s^2 4p^6$; [1]
 Do not accept noble gas shortcut. No subscripts.
- (d) Mg^{2+} ; [1]
- (e) Al^{3+} , O^{2-} , Ne, Na^+ , F^- , N^{3-} ; [2]
 Award [2] for any three, [1] for any two.
3. $n(\text{Fe}_2\text{O}_3) = 30 \times 10^3 \div 159.7 / n(\text{Fe}_2\text{O}_3) = 188 \text{ mol}$;
 $n(\text{C}) = 5.0 \times 10^3 \div 12.01 / n(\text{C}) = 416 \text{ mol}$;
 Fe_2O_3 is the limiting reagent or implicit in calculation;
 $n(\text{Fe}) = 2 \times n(\text{Fe}_2\text{O}_3) = 2 \times 188 = 376 \text{ mol}$;
 $m(\text{Fe}) = 376 \times 55.85 = 21 \text{ kg}$; [5]
 Accept 2sf or 3sf, otherwise use –1(SF).
 Correct final answers score [5].
 Allow ECF.
4. (a) (i) (a species that) gains electrons (from another species) / causes electron loss; [1]
 (ii) changes by 3;
 reduced because its oxidation number decreased $+6 \rightarrow +3$ / $6+ \rightarrow 3+$ / it has gained electrons; [2]
- (b) (i) $\text{C}_6\text{H}_8\text{O}_6 \rightarrow \text{C}_6\text{H}_6\text{O}_6 + 2\text{H}^+ + 2\text{e}^-$; [1]
 (ii) $\text{C}_6\text{H}_8\text{O}_6 + 2\text{Fe}^{3+} \rightarrow \text{C}_6\text{H}_6\text{O}_6 + 2\text{H}^+ + 2\text{Fe}^{2+}$; [1]

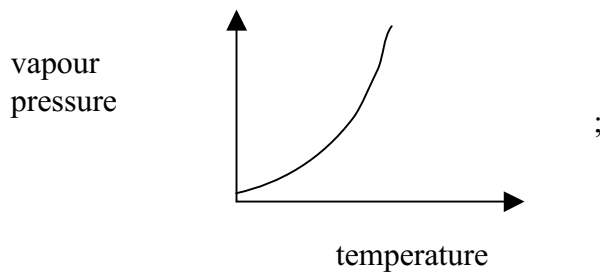
5. (a) same general formula;
 successive members differ by CH_2 ;
Do not allow elements or just “they”.
 similar chemical properties;
Allow same/constant.
 gradual change in physical properties;
Do not allow change periodically.
 same functional group;
Award [1] each for any two. [2 max]
- (b) add bromine (water);
 alkanes – no change / stays or turns brown;
Allow red-brown or any combination of brown, orange or yellow.
 alkenes – bromine (water) decolorizes;
Do not allow clear or discoloured.
- or*
- add (acidified) KMnO_4 ;
 alkanes – no change;
 alkenes – MnO_4^- decolorizes / brown / black; [3]
Do not accept addition of H_2 or HBr .
- (c) butan-1-ol: butanal;
 butanoic acid;
- butan-2-ol: butanone;
- 2 methylpropan-2-ol: no oxidation; [4]
Also accept correct structures. Where both name and structure given structure must be correct and name largely correct.

SECTION B

6. (a) $K / K_c = [\text{SO}_3]^2 \div [\text{SO}_2]^2 [\text{O}_2]$; [1]
Exactly as written.
Accept correct K_p expression.
- (b) (i) vanadium(V) oxide / (di)vanadium pentaoxide / $\text{V}_2\text{O}_5/\text{Pt}$; [1]
Allow just vanadium oxide but not incorrect formula.
- (ii) catalyst does not affect the value of K_c ;
 forward and reverse rate increase equally/by the same factor;
 catalyst increases the rate of the reaction;
 (by providing an alternative path for the reaction with) lower activation energy; [4]
- (c) more energetic collisions / more molecules have energy greater than activation energy;
 more frequent collisions; [2]
Do not accept more collisions without reference to time.
- (d) (i) shifts equilibrium position to the products/right;
 to the side with least gas molecules or moles / lower volume of gas; [2]
- (ii) shifts equilibrium position to the products/right;
 to compensate for loss of SO_3 / produce more SO_3 ; [2]
- (e) exothermic;
 K_c decreases with increasing temperature / back reaction favoured / heat used up /
 OWTTE; [2]
- (f) $n(\text{SO}_2)_{\text{at equilibrium}} = 1.50 - 0.50 = 1.00 \text{ mol}$;
 $n(\text{O}_2)_{\text{at equilibrium}} = 2.00 - 0.250 = 1.75 \text{ mol}$;
- $[\text{SO}_2] = 1.00 \div 1.50 = 0.667 \text{ mol dm}^{-3}$, $[\text{O}_2] = 1.75 \div 1.50 = 1.17 \text{ mol dm}^{-3}$
 $[\text{SO}_3] = 0.500 \div 1.50 = 0.333 \text{ mol dm}^{-3}$;
- $K_c = (0.333)^2 \div 1.17 \times (0.667)^2$;
 $= 0.213 \text{ dm}^3 \text{ mol}^{-1} / 0.214 \text{ dm}^3 \text{ mol}^{-1}$; [5]
Allow ECF.
If $0.202 \text{ dm}^3 \text{ mol}^{-1}$ is given award [4], this is obtained by premature rounding.
Award [5] for correct answer with units.

- (g) (i) the greater the strength of the intermolecular forces the greater the enthalpy of vaporization/*OWTTE*;
 pentane has only van der Waals' forces between molecules;
 propanoic acid has H-bonding (as well as van der Waals' forces); [3]

(ii)

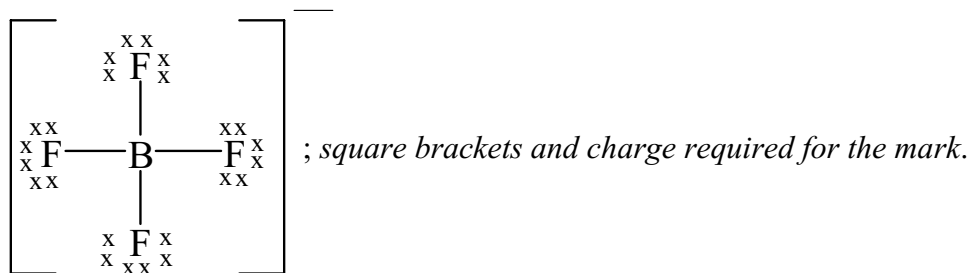
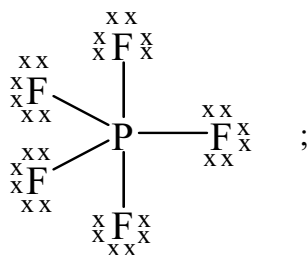
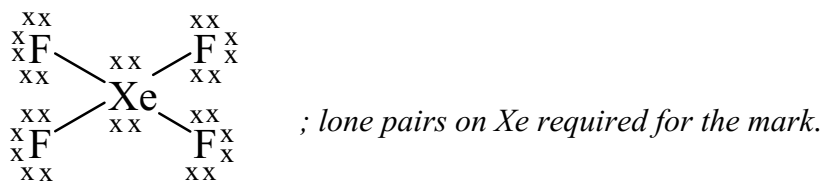


1st mark: graph goes upwards with T;

2nd mark: curve as shown;

as temperature increases (more) molecules have enough energy to overcome intermolecular / attractive forces; [3]

7. (a)



Accept any combination of dots, crosses and lines.
Penalise missing fluorine lone pairs once only.

[3]

(b) XeF_4

Square planar and 90° ;

PF_5

trigonal bipyramid and 90° and 120° ;

BF_4^-

Tetrahedral and $109.5^\circ/109^\circ$;

Allow clear suitable diagrams instead of name.

No ECF from (a).

[3]

(c) hybridization: mixing / merging of atomic orbitals;

N_2 – sp;

N_2H_2 – sp^2 ;

N_2H_4 – sp^3 ;

[4]

(d) σ bonds (result from the) overlapping of orbitals end to end / along inter-nuclear axis;
 π bonds (result from the) overlapping of parallel/sideways p orbitals;

(single bonds) σ bonds only;

(double bonds) have a σ bond and a π bond;

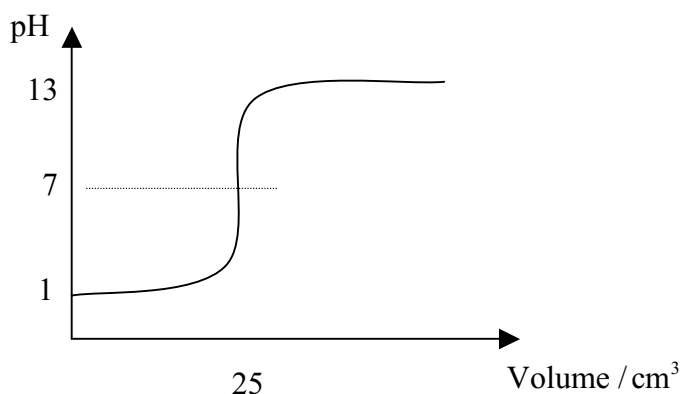
Suitable clear and labelled diagrams acceptable for all marks.

[4]

- (e) (i) electron removed from higher energy level / further from nucleus / greater atomic radius;
increased repulsion by extra inner shell electrons / increased shielding effect; [2]
- (ii) $\text{Mg}^{2+}(\text{g}) \rightarrow \text{Mg}^{3+}(\text{g}) + \text{e}^-$;
(even though) valence electrons in the same shell/main energy level / Mg^{2+} has noble gas configuration;
Mg has greater nuclear/core charge/more protons; [3]
- (f) (i) Mg has twice/more delocalized electrons as Na;
the ionic charge is twice as big/greater in Mg than Na;
sodium ion is larger than magnesium ion;
attraction of ions and electrons is less in sodium/greater in magnesium; [3 max]
Correct discussion of charge density gains 2nd and 3rd mark.
Award [1] each for any three.
- (ii) SO_2 has (weak) intermolecular/van der Waals' force/dipole – dipole;
 MgO has (strong) ionic bonds;
Ionic bonding is stronger than intermolecular attraction (OWTTE); [3]

8. (a) (i) $\text{pH} = -\log[\text{H}^+]$; [1]

- (ii) curve should include the following:
 starting $\text{pH} = 1$;
 equivalence point: 25.0 cm^3 of NaOH ;
 pH at equivalence point = 7;
 pH to finish = 12–13;



Penalise [1] if profile incorrect.

- (iii) $K_a = 10^{-4.76} / 1.74 \times 10^{-5}$;

$$K_a = [\text{H}^+]^2 \div [\text{CH}_3\text{COOH}] / 1.74 \times 10^{-5} = \frac{[\text{H}^+]^2}{0.100};$$

$$[\text{H}^+] = 1.32 \times 10^{-3} \text{ (mol dm}^{-3}\text{)};$$

starting $\text{pH} = 2.88$;

Accept 3sf.

Award [4] for correct pH.

Allow ECF.

pH at equivalence point: 8–9;

- (b) (i) HIn is a weak acid;
 $\text{HIn} \rightleftharpoons \text{H}^+ + \text{In}^-$ and two colours indicated;

In acid equilibrium moves left or vice versa;

- (ii) phenolphthalein / phenol red / bromothymol blue;
 colour change of indicator occurs within the range of pH at equivalence point / on vertical part of graph;

- (c) (i) specific examples of weak base and its salt / specific strong acid and weak base;
Name of structure acceptable.
e.g. NH_3 and NH_4Cl .

- (ii) pH changes very little / most acid neutralized by base;
 equation from (i);
Any other suitable example.
e.g. $\text{NH}_3 + \text{H}^+ \rightarrow \text{NH}_4^+$ / $\text{NH}_4\text{OH} + \text{H}^+ \rightarrow \text{NH}_4^+ + \text{H}_2\text{O}$.

- (d) *Brønsted-Lowry acid*
a proton donor;

Lewis acid
electron pair acceptor;

Brønsted-Lowry acid
Any suitable equation;

Lewis acid – BF_3 / AlCl_3 / transition metal ions that form complex ion with ligands;

For example



Or any suitable equation.

- (e) acidic;
 $[\text{Al}(\text{H}_2\text{O})_6]^{3+}$ is (weak) acid due to the formation of H^+ /



9. (a) (i) CH_2CH_2 ; [1]

(ii) $\begin{array}{c} \text{HOOCCHNH}_2 \\ | \\ \text{CH}_3 \end{array}$; [1]

Allow appropriate acyl chloride.

(iii) $\text{H}_2\text{N}(\text{CH}_2)_6\text{NH}_2$; [2]
 $\text{HOOC}(\text{CH}_2)_4\text{COOH}$;

Allow correct alternative.

Accept correct names as alternatives.

If correct structure and incorrect name given, award the mark.

Penalise COOH – C once only.

(b) (addition polymers) contain $\text{C}=\text{C}$ / $\text{C}\equiv\text{C}$; [2]
 (condensation polymers) contain two reactive/functional groups;

(c) HCOOCH_3 ; [2]
 methyl methanoate;
Accept other correct alternative.

(d) (i) methanol / methyl alcohol; [3]
 heat and acid catalyst / H^+ ;
 $\text{CH}_3\text{OH} + \text{CH}_3\text{COOH} \rightarrow \text{CH}_3\text{COOCH}_3 + \text{H}_2\text{O}$;

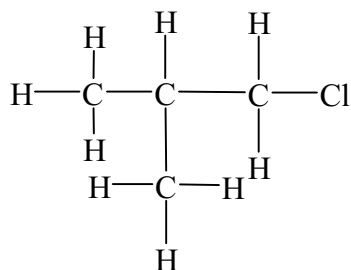
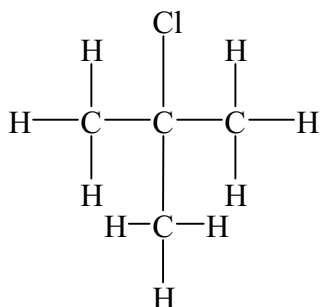
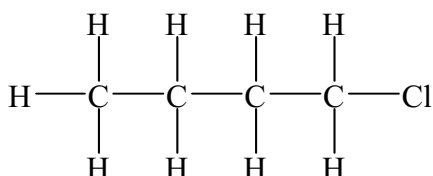
(ii) *physical properties*
 ethanoic acid has a higher boiling point / ester has a lower boiling point;
 ethanoic acid has vinegar smell, ester has sweet/fruit smell;
Must specify one smell.
 ethanoic acid is more soluble in water than methyl ethanoate / methyl ethanoate is more soluble in non-polar solvents than ethanoic acid;
 ethanoic acid (in water) has a $\text{pH} < 7$, ester (in water) has a $\text{pH} = 7$; [2 max]
Award [1] each for any two.

(iii) *ethanoic acid*
 3:1;
methyl ethanoate
 1:1; [2]
Allow 3:3.

- (e) (i) 2 – chlorobutane is the optical isomer;
has a chiral carbon/asymmetric carbon atom / 4 different groups around central atom; [2]

- (ii) pass plane polarized light through (two separate) samples;
each sample will rotate the polarized light in the opposite direction; [2]

- (iii) [2]



*Award [2] marks for 3 and [1] mark for 2 structures.
Penalise missing H atoms once only.*

- (iv) 1-chlorobutane / 1-chloro–2–methylpropane; [1]
Accept structures.

(v) *mechanism*

curly arrow from O of OH^- joined to C, and from C-Cl bond to Cl;

transition state structure with partial bonds to OH and Cl and a negative charge;

product: $\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{OH}$ / $\text{CH}_3\text{CH}(\text{CH}_3)\text{CH}_2\text{OH}$;

[3]

e.g.

